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**Inversion of TES and
MOPITT data**

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Inversion analysis of carbon monoxide emissions using data from the TES and MOPITT satellite instruments

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Abstract

We conduct an inverse modeling analysis of measurements of atmospheric CO from the TES and MOPITT satellite instruments using the GEOS-Chem global chemical transport model. This is the first quantitative analysis of the consistency of the information provided by these two instruments on surface emissions of CO in an inverse modeling context. We focus on observations of CO for November 2004, when the climatological emission inventory in the GEOS-Chem model significantly underestimated the atmospheric abundance of CO as observed by TES and MOPITT. We find that both datasets suggest significantly greater emissions of CO from sub-equatorial Africa and the Indonesian/Australian region. The a posteriori emissions from sub-equatorial Africa based on TES and MOPITT data were 173 Tg CO/yr and 184 Tg CO/yr, respectively, compared to the a priori of 95 Tg CO/yr. In the Indonesian/Australian region, the a posteriori emissions inferred from TES and MOPITT data were 155 Tg CO/yr and 185 Tg CO/yr, respectively, whereas the a priori was 69 Tg CO/yr. The differences between the a posteriori emission estimates obtained from the two datasets are generally less than 20%, and are likely due to the different spatio-temporal sampling of the measurements. The a posteriori emissions significantly improve the simulated distribution of CO, however, large regional residuals remain, reflecting systematic errors in the analysis. For example, the a posteriori emissions obtained from both datasets do not completely reduce the underestimate in the model of CO column abundances over the southern tropical Atlantic, southern Africa, and over the Indian Ocean, where biases of 3–7% remain. Over eastern Asia the a posteriori emissions overestimate the CO column abundances by about 3–6%. These residuals reflect the sensitivity of the top-down source estimates to systematic errors in the analysis. Our results indicate that improving the accuracy of top-down emission estimates will require further characterization of model biases (chemical and transport) and the use of spatial-temporal inversion resolutions consistent with the information content of the observations.

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1 Introduction

Inverse modeling has emerged as a powerful tool for estimating surface emissions of environmentally important trace gases. In this context, inverse modeling of observations of atmospheric CO has become a widely used tool to improve “bottom-up” estimates of surface emissions of CO, which are highly uncertain. Atmospheric CO is a product of incomplete combustion as well as a byproduct of the oxidation of atmospheric hydrocarbons. Accurate and precise estimates of emissions of CO are important since atmospheric CO plays a critical role in determining the oxidative capacity of the atmosphere as the primary sink of OH, and it is also a useful tracer of atmospheric transport.

As a combustion product, atmospheric CO provides a useful proxy for combustion-related emissions of other precursors of tropospheric O₃. It is in this context that we conduct an inverse modeling analysis, using the GEOS-Chem chemical transport model (CTM), to quantify emissions of CO for fall 2004. We focus on November 2004 when observations from the Tropospheric Emission Spectrometer (TES) and the Measurement of Pollution in The Troposphere (MOPITT) satellite instruments revealed significantly higher abundances of atmospheric CO in the southern tropics than simulated by the GEOS-Chem model. In a companion paper by Bowman et al. (2007)¹ a detailed analysis is presented of the impact on tropical tropospheric O₃ of the changes in emissions of CO (and the implied changes in emissions of other O₃ precursors) inferred from the inversion analysis conducted here.

The dominant source of variability in atmospheric CO in the tropics is associated with biomass burning. Accurately quantifying emissions of CO from these sources is

¹Bowman, K. W., Jones, D. B. A., Logan, J. A., Worden, H., Boersma, F., Kulawik, S., Chang, R., Osterman, G., and Worden, J.: Impact of surface emissions to the zonal variability of tropical tropospheric ozone and carbon monoxide for November 2004, Atmos. Chem. Phys. Discuss., submitted, available from: <http://www.atmosphysics.utoronto.ca/~jones/publications.html>, 2007.

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particularly challenging as they exhibit significant spatial and temporal variability and are sensitive to variations in the climate system such as the El Niño Southern Oscillation (ENSO) (Duncan et al., 2003; Van der Werf et al., 2004). For example, van der Werf et al. (2006) found that the standard deviation of the variability in the bottom-up estimates for carbon emissions biomass burning in Indonesia for 1997–2004 was 1.3 times as large as the mean emissions for the period. In late 2004 there was a weak El Niño in the tropical Pacific, which could account for some of the discrepancy between the model and observations. In our analysis we focus on quantifying the total combustion-related source of CO from different continental regions, with the assumptions that biomass burning is the dominant source of direct emissions of CO in the tropics and that interannual variability in biomass burning contributes largely to the discrepancy between the observations and the modeled CO distribution obtained with the climatological emission inventory.

In the past decade several inverse modelling studies of CO have been conducted using surface, aircraft, and satellite measurements (e.g. Bergamaschi et al., 2000a, b; Kasibhatla et al., 2002; Pétron et al., 2002, 2004; Palmer et al., 2003; Heald et al., 2004; Arellano et al., 2004, 2006; Müller and Stavrakou, 2005; Stavrakou and Müller, 2006; Kopacz et al., 2007). These studies have all produced different “top-down” estimates of the regional sources of CO (see discussion in Duncan et al., 2007), reflecting differences in inversion frameworks, atmospheric models (e.g. Arellano and Hess, 2006), and differences in the datasets employed in the analyses. A particular challenge for the earlier inversion analyses, which employed surface or aircraft measurements, was the limited spatial and temporal coverage provided by the observations. The recent satellite measurements of atmospheric CO, on the other hand, offer significantly greater spatio-temporal coverage and therefore provide more reliable constraints on regional surface emissions of CO (Heald et al., 2004). The MOPITT instrument, launched in December 1999, provided the first continuous global measurements of atmospheric CO. More recently, space-based measurements of CO have become available from the TES instrument, launched in July 2004. A second objective of this study is to evaluate

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the consistency of the top-down constraints on surface emissions of CO provided by observations from these two satellite instruments. We also examine the impact on the source estimates of differences in the spatio-temporal sampling of the instruments.

The broad range of published top-down estimates of regional CO emissions reflects the fact that Bayesian inverse modeling is sensitive to systematic errors in the inverse models. Another objective of this study is to demonstrate that, despite the significantly increased spatio-temporal coverage offered by data from TES and MOPITT, the CO source estimates inferred from these datasets are sensitive to the presence of systematic errors, such as those associated with the aggregation of the emission regions or the neglect of the non-linearity of the atmospheric chemistry of CO. Assuming that the chemistry of CO is linear is a potential source of systematic error in most inverse modeling analyses of atmospheric CO, which is not well addressed in the literature. Inversion studies typically prescribe abundances of atmospheric OH to account for the chemical sink of CO. However, if large changes in emissions of CO, such as those associated with enhanced biomass burning, are required in an inverse model to accommodate observations of CO, then there should be a concomitant increase in the model of emissions of O₃ precursors such as nitrogen oxides (NO_x), methane (CH₄), and non-methane hydrocarbons (NMHC), which will, in turn, perturb the atmospheric abundance of OH. We examine here the potential feedback of perturbations in the atmospheric abundance of OH associated with changes in biomass burning on the atmospheric abundance of CO.

In Sect. 2 we begin with a brief description of the CO profile retrievals from the TES and MOPITT instruments. We describe the GEOS-Chem model in Sect. 3. The inversion methodology is presented in Sect. 4, followed by a discussion of the inversion results in Sect. 5. In Sect. 6 we provide a summary of the results and a discussion of their implications for future inversion analyses of atmospheric CO.

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2 The MOPITT and TES Instruments

2.1 MOPITT

The MOPITT instrument (Drummond and Mand, 1996) was launched on 18 December 1999 on NASA's Terra spacecraft in a sun-synchronous polar orbit at an altitude of 705 km with an equator crossing time of 10:30 a.m. local time. It is gas correlation radiometer that measures thermal emission in the $4.7\ \mu\text{m}$ region of the spectrum and has a spatial resolution of $22\text{ km}\times 22\text{ km}$. The MOPITT observation strategy consists of a 612 km cross-track scan that provides high data density; the instrument achieves nearly complete global coverage every 3 days.

Vertical profiles of CO are retrieved from the radiance measurements using an optimal estimation approach, described by Deeter et al. (2003). The retrieved mixing ratios are reported on 7 altitude levels, from the surface to 150 hPa. The data have been validated by inter-comparison with aircraft and other in-situ measurements (Emmons et al., 2004, 2007). In our analysis we employ version 3 MOPITT data and use profiles between 700–250 hPa, as these are the levels with most of the information in the retrievals (Emmons et al., 2007). The retrieved profiles can be expressed as a linear estimate of the true atmospheric state

$$\hat{\mathbf{x}}^{\text{MOP}} = \mathbf{x}_a^{\text{MOP}} + \mathbf{A}^{\text{MOP}}(\mathbf{x}^{\text{true}} - \mathbf{x}_a^{\text{MOP}}) + \boldsymbol{\varepsilon} \quad (1)$$

where $\mathbf{x}_a^{\text{MOP}}$ is the MOPITT a priori CO profile, \mathbf{x}^{true} is the true atmospheric state vector, \mathbf{A}^{MOP} is the MOPITT averaging kernel matrix, and $\boldsymbol{\varepsilon}$ is the measurement error. The averaging kernel is given by $\mathbf{A} = \frac{\partial \hat{\mathbf{x}}^{\text{MOP}}}{\partial \mathbf{x}^{\text{true}}}$ and represents the sensitivity of the MOPITT retrieval to the true state of the atmosphere and provides a measure of the vertical resolution of the retrieval.

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2.2 TES

The TES instrument is a high resolution Fourier transform spectrometer that was launched on the Aura spacecraft on 15 July 2004. It measures thermal emission between 3.3–15.4 μm in both nadir and limb modes (Beer et al., 2001). The satellite is in a sun-synchronous orbit at an altitude of 705 km with an inclination of 98.2° and a repeat cycle of 16 days. The instrument operates in two observational modes: a global survey mode in which the observations are spaced about 5° along the orbit track, and a step-and-stare mode in which the observation are made every 40 km long the orbit. The horizontal resolution of the nadir observations is 8 km×5 km. Similar to MOPITT, vertical profiles of CO are retrieved from radiance measurements in the 4.7 μm spectral region using an optimal estimation approach. The TES retrievals, however, are performed for the logarithm of the mixing ratio of CO. A detailed discussion of the TES retrievals is given in Bowman et al. (2006). As for MOPITT, the TES retrievals can be expressed as a linear representation of the true atmospheric state

$$\hat{\mathbf{x}}^{\text{TES}} = \mathbf{x}_a^{\text{TES}} + \mathbf{A}^{\text{TES}}(\mathbf{x}^{\text{true}} - \mathbf{x}_a^{\text{TES}}) + \boldsymbol{\varepsilon} \quad (2)$$

where $\mathbf{x}_a^{\text{TES}}$ is the TES a priori CO profile, \mathbf{A}^{TES} is the TES averaging kernel matrix, and $\boldsymbol{\varepsilon}$ is the measurement error. For consistency with the inversion using MOPITT data, we restrict the TES profiles ingested in the inversion to between 800–250 hPa.

We employ 6 global surveys of TES CO data that were obtained during 4–16 November 2004. There were a limited number of global survey data available before and after this period in fall 2004. We use version V001 of the TES data as the major change in V002 was associated with improvements in the calibration algorithms to reduce biases in the TES radiances, which decreased the bias in the TES O₃ retrievals in the upper troposphere (Worden et al., 2007). The most significant change in the TES CO product occurred as a result of the warm-up of the optical bench in the instrument in December 2005 to correct for decreasing signal strength in the 1A1 filter. Before warm-up the sensitivity of the TES CO retrievals had dropped to less than 1 degree of freedom for signal (DOFS) compared to a sensitivity of 1–2 DOFS after launch and after the

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warm-up. The DOFS is the trace of the averaging kernel matrix and provides a measure of the number of independent pieces of information on the vertical distribution of CO available in the retrieved profile. It is, in part, because of the low DOFS in the CO retrievals in fall of 2005 that we restrict our analysis to fall 2004.

5 A detailed comparison of the TES and MOPITT profile retrievals of CO was presented by Luo et al., (2007a), while validation of the TES CO retrievals with aircraft observations was conducted by Luo et al. (2007b). Luo et al. (2007a) found that despite the differences in the measurement and retrieval techniques of TES and MOPITT, both datasets provide about 0.5–2 degrees of freedom for signal (DOFs) in the troposphere.
 10 Luo et al. (2007a) found that after accounting for the a priori profiles incorporated in the retrievals and differences in the averaging kernels between the instruments the two datasets were in good agreement, with an absolute mean difference of less than 5% in the column abundances of CO. In the vertical profiles, the largest mean differences between the two datasets were –4.8% at 150 hPa (for TES compared to MOPITT),
 15 whereas the smallest mean differences were –0.2% at 850 hPa (Luo et al., 2007a). The focus of the work presented here is to assess if this agreement between the two datasets imply consistency in the constraints that they provide on surface emissions of CO when the data are incorporated in an inverse model.

3 Inversion methodology

20 The inversion framework employed here is described in Jones et al. (2003). We obtain optimized estimates of the sources of CO by minimizing the cost function (Rodgers, 2000)

$$J(\mathbf{y}) = (\hat{\mathbf{x}} - F(\mathbf{y}))^T \mathbf{S}_e^{-1} (\hat{\mathbf{x}} - F(\mathbf{y})) + (\mathbf{y} - \mathbf{y}_a)^T \mathbf{S}_a^{-1} (\mathbf{y} - \mathbf{y}_a) \quad (3)$$

25 where $\hat{\mathbf{x}}$ is the observation vector that consists of the retrieved vertical profiles of CO (defined in Eqs. 1 and 2) for MOPITT and TES, respectively), \mathbf{y} is the state vector with elements representing monthly mean emissions from the source regions shown

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in Fig. 1, and \mathbf{y}_a is the a priori state vector, which is based on the emission inventory described in Sect. 4 and is given in Table 1. Note, for consistency with the description of the satellite retrievals in Eqs. (1) and (2), we denote the state vector in the source inversion as \mathbf{y} although the standard optimal estimation nomenclature as described in Rodgers (2000) uses \mathbf{x} as the state vector. \mathbf{S}_ε is the error covariance matrix of the observation vector and \mathbf{S}_a is the a priori covariance matrix. The forward model $F(\mathbf{y})$ reflects the transport of the CO emissions in the GEOS-Chem model, and accounts for the vertical sensitivity of TES and MOPITT and the a priori CO profile used in the retrievals in the two datasets. It is given by

$$F^{\text{MOP}}(\mathbf{y}) = \mathbf{x}_a^{\text{MOP}} + \mathbf{A}^{\text{MOP}}(H(\mathbf{y}) - \mathbf{x}_a^{\text{MOP}}) \quad (4)$$

and

$$F^{\text{TES}}(\mathbf{y}) = \mathbf{x}_a^{\text{TES}} + \mathbf{A}^{\text{TES}}(\ln[H(\mathbf{y})] - \mathbf{x}_a^{\text{TES}}) \quad (5)$$

where $H(\mathbf{y})$ is the GEOS-Chem model which transports and chemically transforms the CO emissions (\mathbf{y}).

We assume Gaussian error statistics for the observation and a priori errors, which yields the following maximum a posteriori (MAP) solution for the minimization of the cost function (Rodgers, 2000),

$$\mathbf{y}_{i+1} = \mathbf{y}_i + (\mathbf{K}_i^T \mathbf{S}_\varepsilon^{-1} \mathbf{K}_i + \mathbf{S}_a^{-1})^{-1} [\mathbf{K}_i^T \mathbf{S}_\varepsilon^{-1} (\hat{\mathbf{x}} - F(\mathbf{y}_i)) - \mathbf{S}_a^{-1} (\mathbf{y}_i - \mathbf{y}_a)]. \quad (6)$$

Here \mathbf{y}_i and $\mathbf{K}_i = \partial F(\mathbf{y}_i) / \partial \mathbf{y}$ are estimates of the state vector and the Jacobian matrix, respectively, at the i th iteration. It is necessary to use an iterative approach in solving for the MAP solution because of the slight nonlinearity introduced by the logarithm in the TES forward model (Eq. 5). We obtain the solution in Eq. (6) using a Gauss-Newton method (Rodgers, 2000), with the sequential approach outlined in Jones et al. (2003). The Jacobian is estimated using separate atmospheric tracers of CO for emissions from each region in the state vector, as described in Sect. 4. In each region we solve for the total emissions of CO.

In constructing the a priori error covariance matrix we assume that emissions from the different sources are uncorrelated and have a uniform uncertainty of 50%, following Palmer et al. (2003), except for emissions from North American and Europe for which Palmer et al. (2003) assumed an uncertainty of 30%. We specify a uniform a priori constraint to more clearly assess the impact of the two datasets on the source estimates.

The observation error covariance matrix consists of the retrieval error covariance, the model error covariance, and the representativeness error covariance. Previous inverse modeling studies (Palmer et al., 2003; Jones et al., 2003; and Heald et al., 2004) for atmospheric CO have shown that the model error dominates the observation error covariance, with the representativeness and measurement errors providing a small contribution to the total error. In their inversion analysis of Asian emissions of CO, Heald et al. (2004) examined the sensitivity of the estimates of the CO sources to the specification of the model error. They compared the impact on the source estimates of specifying a uniform model error with that obtained using the complete model error covariance structure for the GEOS-Chem simulation of CO, as determined by Jones et al. (2003). They found that there was sufficient information in the MOPITT data such that the source estimates for regions with strong biomass burning emissions of CO, such as southeast Asia, were insensitive to the specification of the error covariance structure. Assuming a uniform estimate of 20% for the total observation error in their analysis, for example, did not significantly influence the estimates for source regions such as southeast Asia. In contrast, they found that the weaker regional sources, such as estimates for emissions from western China and Japan and Korea were influenced strongly by the choice of error covariance. As our focus here is on the dominant, continental-scale, biomass burning signals in the tropics, we assume a uniform observation error of 20%. This approach is similar to previous inverse modeling studies of CO such as Kasibhatla et al. (2002) and Pétron et al. (2002). We also neglect the correlations in the model error, but account for the influence of the averaging kernels from the instruments on the model error, which captures the vertical correlations associated

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with the vertical smoothing of the retrievals.

4 The GEOS-Chem model

The GEOS-Chem model (Bey et al., 2001) is a global three-dimensional CTM driven by assimilated meteorological observations from the NASA Goddard Earth Observing System (GEOS-4) from the Global Modeling and data Assimilation Office (GMAO). The meteorological fields have a horizontal resolution of $1^\circ \times 1.25^\circ$ with 55 levels in the vertical, and a temporal resolution of 6 h (3 h for surface fields). We employ version 7-02-04 of GEOS-Chem (<http://www-as.harvard.edu/chemistry/trop/geos>), with the resolution of the meteorological fields degraded to 2° latitude \times 2.5° longitude. Recent applications of the model have been described in a range of studies (e.g. Suntharalingam et al., 2004; Heald et al., 2004; Park et al., 2005; Hudman et al., 2007; and Wang et al., 2007). The model includes a complete description of tropospheric O_3 - NO_x -hydrocarbon chemistry, including the radiative and heterogeneous effects of aerosols.

The GEOS-Chem simulation of CO for 4–15 November 2004 is shown in Fig. 2. The model significantly underestimates the CO abundances as observed by TES and MOPITT, reflecting the climatological emission inventories used in the simulation. Anthropogenic emissions of CO in this version of GEOS-Chem are as described by Duncan et al. (2007), with emissions for the base year of 1985 in the inventory scaled to 1998. For biomass burning we employ the emission inventory of Duncan et al. (2003), whereas for biofuel combustion we use estimates from Yevich and Logan (2003). The total a priori global emissions of CO, as well as the total secondary source of CO from the oxidation of methane and NMHC is given in Table 1.

In our inversion analysis we specify monthly mean concentrations of OH to linearize the chemistry of CO. This approach has been used previously for the inverse modeling of CO using the GEOS-Chem model (e.g. Palmer et al., 2003, 2006; Jones et al., 2003; Heald et al., 2004; and Kopacz et al., 2007). Linearization of the chemistry enables us to efficiently calculate the Jacobian in the inversion analysis by specifying a separate

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tracer for each region considered in the state vector. These OH fields were obtained from an earlier version of the full chemistry simulation of GEOS-Chem (Evans and Jacob, 2005).

5 Results

5 The results of the inversion analysis using data from TES and MOPITT for early November 2004 are shown in Fig. 3 and Table 1. They suggest significantly greater emissions of CO in sub-equatorial Africa and the Indonesian/Australian region than the a priori. In sub-equatorial Africa, where the a priori estimate is 95 Tg CO/yr (Table 1), the inferred emission estimate from TES data is 173 Tg CO/yr, while the estimate based
10 on MOPITT data is 184 Tg CO/yr. In the Indonesian/Australian region the a posteriori emission estimates are 155 Tg CO/yr and 185 Tg CO/yr, based on the TES and MOPITT data, respectively, compared to the a priori of 69 Tg CO/yr. Following Palmer et al. (2003) and Heald et al. (2004) we assume here that the seasonality of the sources in the model is correct and, therefore, report the source estimates as annual means al-
15 though the inversion provides constraints on the source estimates only for the October to November period, reflecting the atmospheric lifetime of CO. In the analysis we do not attempt to discriminate between emissions of CO from different source types, such as biomass burning or fuel combustion, however, we assume that the increases in emis-
sions suggested for the tropical regions are associated mainly with greater emissions of CO from biomass burning. The globally averaged source of CO estimated from the
20 TES and MOPITT measurements are 2646 Tg CO/yr and 2761 Tg CO/yr, respectively, compared to the a priori of 2243 Tg CO/yr (Table 1). The agreement between the global estimates from TES and MOPITT is encouraging, but it should be noted that the pre-
scribed OH fields have a global mean OH abundance of $11.3 \times 10^5 \text{ cm}^{-3}$, and biases in
25 the chemistry, as reflected in the OH abundances, would adversely impact the accuracy of the source estimates from both instruments, despite their apparent consistency.

The a posteriori estimates of the CO emissions for sub-equatorial Africa and the

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Indonesian/Australian region based on the TES data thus represent an increase in emissions by a factor of 1.8 and 2.3, respectively, above the a priori values. The large increase in emissions from these regions are consistent with the results of Arellano et al. (2006), who conducted an inversion analysis of MOPITT data for 2000–2001 and reported similarly greater top-down estimates of CO emissions. Arellano et al. (2006) estimated total a posteriori emissions from Indonesia and Oceania (which includes Australia) of about 165 Tg CO/yr, which is comparable to the estimates of 155–185 Tg CO/yr obtained in this study. For sub-equatorial Africa they reported an estimate of 203 Tg CO/yr, slightly larger than the 175–185 Tg CO/yr obtained here. It should be noted that in their analysis, Arellano et al. (2006) quantified separately CO emissions from fuel combustion and biomass burning and found that fuel combustion provided a significant contribution to the total a posteriori emission estimates for Indonesia and sub-equatorial Africa. For example, their a posteriori estimate for emissions from fuel combustion from Indonesia and Oceania were 84 Tg CO/yr and about 5 Tg CO/yr, respectively. In contrast, their a posteriori estimate for biomass burning from these regions was 76 Tg CO/yr. We do not believe that our inversion approach can reliably discriminate between emissions of CO from fuel combustion and biomass burning. However, the consistency between our results for 2004 and those of Arellano et al. (2006) for 2000 does suggest that fuel combustion may indeed be responsible for a large fraction of the emissions as these sources have less interannual variability.

In a companion paper by Bowman et al. (2007)¹ the a posteriori emissions from the inversion are evaluated in the context of their impact on the modeled O₃ distribution. Using a forward model simulation of GEOS-Chem with the O₃ precursor emissions from fuel combustion and biomass burning scaled by the regional scaling factors obtained in the CO inversion, Bowman et al. (2007) showed that the a posteriori emissions provide an improved simulation of O₃ over Indonesia and Australia. Throughout the free troposphere over Indonesia and Australia O₃ increased in the model by as much as 10 ppb, reducing the maximum bias in the modeled O₃ distribution relative to TES by about 40%. In contrast, in the free troposphere over the tropical southern

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Atlantic, Bowman et al. (2007)¹ found that the improvement in the modeled O₃ with the a posteriori emissions was modest, despite the large increases in surface emissions from sub-equatorial Africa, reflecting the greater influence of NO_x from lightning on the budget of O₃ in this region.

The results obtained here show that TES and MOPITT data provide consistent constraints on surface emissions of atmospheric CO. As shown in Table 1, the absolute differences in the source estimates inferred from the two datasets are about 20% or less, with the exception of emissions from North America, which is discussed further below. These differences represent the potential influence on the source estimates of the different spatio-temporal sampling of the TES and MOPITT measurements when the data are incorporated into a regional Bayesian inverse analysis. It is likely that a higher resolution inverse model than that used in this study, which optimizes the emissions at the spatio-temporal resolution of the data, may result in smaller differences between the source estimates inferred from the two datasets.

We conducted a sensitivity analysis using observations from MOPITT for the whole month of November (5–28) to determine the potential impact on the source estimates of using only two week of data on early November. We found that the MOPITT data for 5–28 November data produced results similar to those obtained with data for the first half of November, suggesting that the information from early November is representative of the sources of CO for the October-November period. This is consistent with the simulation experiments of Jones et al. (2003) that suggested two weeks of TES would provide sufficient information to quantify large-scale continental sources of CO.

The inversion analysis can independently quantify emissions from the three continental regions in the southern tropics because transport patterns from these regions are broad and are relatively distinct, reflecting the local meteorology. The distribution of the tagged CO tracers emitted from South America, subequatorial Africa, and the Indonesian/Australian region are shown in Fig. 4. Emissions from South America are entrained into the subtropics in the southern tropical Atlantic and are transported eastward in the westerlies in the subtropics and extratropics of the southern hemisphere.

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Emissions from sub-equatorial Africa, on the other hand, are exported across the Indian Ocean and Australia. Over Australia and Indonesia the dominant contribution to the total abundance of CO in the free troposphere is from local emissions convectively transported out of the boundary layer. In general, over each continental region, convective transport of surface emissions to the upper troposphere and subsequent eastward outflow provides a strong signal for CO in the middle troposphere where TES and MOPITT are most sensitive.

While the focus on this analysis is on the tropics, the inverse model is global in scope and the major discrepancy between the results obtained with TES data and those from MOPITT data is in the estimate for North American emissions. The TES data suggest a significantly reduced North American source of CO compared to the a priori, whereas the MOPITT data suggest a slightly larger source (Table 1). This reduction of the estimate of the North American source with the TES data reflects the fact that during boreal fall North American emissions provide a small contribution to the total CO abundance in the free troposphere at midlatitudes. As shown in Fig. 5, North American emissions account for less than about 16% of the total CO in the middle troposphere at midlatitudes, which means that the signal for North American emissions is more challenging to discriminate from the background, given the noise level in the inversion analysis. We found that with TES data the North American source estimate is correlated with the European estimate ($r=-0.6$) and with the background source of CO ($r=-0.5$). In contrast, with MOPITT data the North American and European source estimates are uncorrelated ($r=-0.09$), while the North American estimate is correlated with the background ($r=-0.6$). This suggests that the estimate for the North American source obtained with both datasets for this period will be sensitive to errors in the background source.

The inversion is less sensitive to the North American source because these emissions are widely distributed at high latitudes in the Arctic region, but in our inversion analysis we neglect retrievals poleward of 60° from both satellites as they tend to be less reliable. At midlatitudes there are episodic transport events, such as warm

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conveyor belts associated with the passage of cold fronts, which transport air out of the North American boundary layer and produce enhanced levels of CO in the free troposphere. These signals, however, are localized spatially and temporally and are therefore not captured by the TES data. MOPITT, on the other hand, because of its greater spatio-temporal sampling density can resolve these synoptic structures (Liu et al., 2006) and therefore provide more constraints on North American emissions.

The discrepancy between the North American estimates illustrates the importance of properly selecting the state vector in the inversion analysis that is consistent with the spatio-temporal sampling of the observation used in the analysis. Ideally, the estimates for those elements of the state vector for which the observations provide little information should remain at the values specified by the a priori. The low estimate of the North American source obtained with the TES data indicate the presence of systematic errors in the inverse model.

The results obtained using both TES and MOPITT data suggest significantly larger emissions from Asia, 511 Tg CO/yr and 531 Tg CO/yr, respectively, compared to the a priori estimate of 367 Tg O/yr. In contrast, the inverse modeling studies of Heald et al. (2004) and Arellano et al. (2006), which inferred CO emissions from MOPITT data, reported estimates for the Asian emissions of CO (excluding emissions from Indonesia) of 282 Tg CO/yr and 402 TgCO/yr, respectively. The differences between the a posteriori estimates of the Asian sources reported by these studies are due, in part, to the fact that these studies used different model configurations. Furthermore, the analyses were focused on different periods: our analysis was carried out for November 2004, whereas the Heald et al. (2004) study was focused on February–April 2001, and Arellano et al. (2006) conducted a time dependent inversion analysis for April 2000–April 2001. These studies will, therefore, be impacted differently by the spatio-temporal sampling of the data, aggregation of the source regions, and by systematic errors associated with the transport and chemistry. It is also possible that some fraction of the differences between the Asian estimates reported here and those from the earlier studies could reflect actual increases in emissions of CO in Asia since 2000. However,

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examination of the residuals from the CO simulation with the a posteriori emissions (discussed below), show that the Asian estimates reported here do provide an overestimate of the Asian sources of CO, and are likely due to systematic errors in the inversion.

As mentioned above, the source estimates obtained from two weeks of MOPITT data are consistent with those inferred from data for the whole month of November 2004. The exception is for north equatorial Africa, for which the inversion using the latter dataset suggests a much larger estimate for the CO emissions compared to results with the former. Biomass burning, which is the dominant source of CO from north equatorial Africa, increases during boreal fall and winter, reaching a maximum in December–January (Duncan et al., 2003). Indeed, fire-counts inferred from the MODIS instrument show more widespread burning in late November 2004, compared to early November. This increased burning later in the month is reflected in the larger source estimate obtained when MOPITT data from late November is included in the inversion analysis.

5.1 Comparison of GEOS-Chem with a posteriori emissions to TES and MOPITT

The a posteriori emissions provide a significantly improved simulation of the distribution of CO, as shown in Table 2 and Figs. 6 and 7. The global mean bias (averaged 60° S–60° N) in the modeled column abundances of CO with respect to the TES data is reduced from –12% to 0.1%, while the bias with respect to the MOPITT data is reduced from –22% to –0.8%. Despite the significant improvement in the global mean CO, the model simulation with the optimized emissions produces large regional biases, as shown in Figs. 6, 7, and Table 2. The residuals of the model fit to the TES and MOPITT data shown in Figs. 6b and 7b and Table 2 indicate that the inversion with both datasets provides an underestimate of 3–7% of the CO column abundances over the southern tropical Atlantic, southern Africa, and over the Indian Ocean.

The optimized emissions inferred from MOPITT data produce an underestimate of CO abundances over the Sahara, the Middle East, and over the North Pacific, while they result in an overestimate of CO abundances across the tropical Pacific and over

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tropical western Africa. These regional biases, however, are compensatory such that the mean bias across the tropics and midlatitudes of the southern hemisphere (0° – 45° S) is small, -3% and -1% for emissions inferred from TES and MOPITT data, respectively. Similarly, in the extratropics of the northern hemisphere (25° – 60° N) the mean biases in the model, relative to the TES and MOPITT data, are reduced from -7% and -18% , respectively, to 0.9% and -2.7% with the a posteriori emissions. The fact that the reduced North American emissions obtained with the TES data do not contribute to a noticeable bias between the model simulation with the a posteriori emissions and the TES observations over North America is an indication that, as discussed above, North American emissions provide a small contribution to the total CO in the free troposphere in boreal fall.

The large a posteriori estimate for Asian emissions represents an overestimate of the Asian sources, as reflected in the large negative residuals over East Asia (Figs. 6b and 7b and Table 2). As mentioned above, this is likely due to a combination of aggregation errors in the inversion analysis and bias in the model transport or chemical fields. With the a priori emissions, the model underestimates the CO abundance over the north Pacific compared to the observations from TES and MOPITT. Emissions of CO from southeast Asia represent a large contribution to the total CO in the upper troposphere over the Pacific. The a priori bias over the North Pacific could reflect either an underestimate in the magnitude of these emissions or a bias in the rate at which these emissions are transported to the upper troposphere (mostly likely by convective transport). By aggregating all of the Asian emissions into one region, the inversion analysis scales all of the Asian emissions in trying to compensate for this underestimate of CO over the North Pacific, potentially resulting in an overestimate of the eastern Asian emissions. As demonstrated recently by Kopacz et al. (2007), conducting the inversion at the resolution of the model would provide maximum flexibility in adjusting the emissions to best reproduce the observations (given the uncertainty of the observations and the model simulation), while minimizing the aggregation errors.

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5.2 Feedback of changes in atmospheric OH on CO

In the inversion analysis we linearized the chemistry of CO by imposing monthly mean concentrations of OH, the main sink for atmospheric CO. This, however, introduces biases in the inversion as observations of CO ingested in the analysis will also reflect the influence of OH concentrations characteristic of different background chemical conditions in the atmosphere. For example, enhanced combustion-related emissions of CO in Indonesia/Australia, as suggested by the observations, will result in increased emissions of O₃ precursors, leading to elevated atmospheric abundances of O₃ and OH, and consequently to suppressed concentrations of CO. To assess the magnitude of this feedback on the atmospheric concentrations of CO we conducted a forward model simulation with the full nonlinear chemistry and with the combustion-related emissions of NO_x (from biomass burning and biofuel and fossil fuel combustion) scaled uniformly according to the regional scaling factors obtained in the inversion analysis of the CO data. We scale only NO_x emissions in the simulation and compare the resulting CO distribution with that obtained with the a priori NO_x emissions. We neglect possible errors associated with assuming that the relative contribution of different source types to the total emissions of NO_x is the same as for emissions of CO in these regions.

The change in the abundance of CO obtained with the scaled NO_x emissions is shown in Fig. 8. The increased emissions of NO_x from Indonesia and Australia results in a reduction of CO by as much as 7–10 ppb, corresponding to about 10% of the total CO abundance. This decrease in CO is a result of increased tropospheric O₃, and thus OH, in the Indonesian/Australian region. Higher concentrations of NO_x produce an increase in O₃ throughout the southern tropics, with the largest increases, of about 35%, in the middle/upper troposphere over Indonesian/Australian (not shown). A decrease of about 7–10 ppb in CO over Indonesia and Australia represents about 20–30% of the contribution of emissions from this region to the total abundance of CO (Fig. 5) and suggests that neglecting the chemical feedback of changes in surface emissions on the abundance of OH, and thus CO, could introduce biases in the a posteriori estimates of

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the sources of CO. Indeed, in their inversion analysis of surface measurements of CO and GOME observations of NO₂, Müller and Stavrakou (2005) found that their a posteriori CO emissions obtained by simultaneously optimizing the CO and NO_x sources provided a better simulation of aircraft observations of CO than those estimated from only the surface CO data.

6 Summary and discussion

We have conducted an inverse modeling analysis of observations of atmospheric CO from the TES and MOPITT satellite instruments to assess the constraints that these data provide on estimates of surface emissions of CO. We focused our analysis on observations from November 2004, during the biomass burning season in the southern hemisphere, as emissions of CO from biomass burning represent the dominant source of variability in atmospheric CO in the tropics. Observations from 4–15 November 2004, indicated that the climatological emissions inventory in the GEOS-Chem model significantly underestimated the abundance of CO observed by both instruments. We used a maximum a posteriori inverse modeling approach to quantify the magnitude of emissions of CO most consistent with the observations. Although our focus was on the tropics, the inversion analysis was global, employing profile retrievals of CO from MOPITT and TES between 60° S–60° N.

The TES and MOPITT data provided consistent information on the CO sources; differences between the a posteriori emission estimates obtained from the two datasets were generally less than 20%. We found that both datasets suggested significantly greater emissions of CO (by a factor of 2–3) from sub-equatorial Africa and from the Indonesian/Australian region in November 2004. The a posteriori emissions from sub-equatorial Africa based on TES and MOPITT data were 173 Tg CO/yr and 184 Tg CO/yr, respectively, compared to the a priori of 95 Tg CO/yr. In the Indonesian/Australian region, a posteriori emissions inferred from TES and MOPITT data were 155 Tg CO/yr and 185 Tg CO/yr, respectively, whereas the a priori was 69 Tg CO/yr. In

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contrast, the a posteriori emissions from South America were not significantly different from the a priori.

We found that while the source estimates for the TES and MOPITT were consistent, the inversion using both datasets was sensitive to the presence of systematic errors in the analysis. The inversion produced much larger estimates for emissions from Asia, 511 Tg CO/yr based on TES data and 531 Tg CO/yr based on MOPITT data, which are greater than most previously published estimates of Asian emissions. These a posteriori emissions result in large residuals over East Asia, with the model simulation providing an overestimate of the observed CO from both TES and MOPITT. In contrast, the model simulation with the a priori emissions underestimated the observed CO over this region. The residual bias over East Asia is likely a result of aggregation errors and biases in the model transport. The contribution of aggregation error to the residual bias could be minimized by constraining the Asian emissions at a higher spatial resolution thereby allowing the inverse model to adjust the Asian emissions locally rather than uniformly scaling all of the emissions across Asia. Ideally, the inversion should be conducted at the resolution of the model using an adjoint modeling approach such as that employed by Stavrakou and Müller (2006) and Kopacz et al. (2007).

We also examined the feedback on atmospheric CO of variations in tropospheric OH associated with changes in biomass burning emissions, as suggested by the a posteriori CO source estimates. Using a forward model simulation of the full tropospheric chemistry with NO_x emissions from combustion sources scaled uniformly based on the regional scaling factors inferred in the CO inversion analysis produced increases in O₃ and OH throughout the southern tropics with the largest increases over Indonesia/Australia. The abundance of O₃ increased by about 35% in the middle/upper troposphere over Indonesian/Australia. Bowman et al. (2007)¹ show that the regional O₃ response to enhanced combustion-related emissions is more complex over regions such as South America and sub-equatorial Africa when emissions of hydrocarbons such as acetaldehyde, acetone, and formaldehyde are considered in addition to emissions of NO_x. In response to the changes in O₃ and OH, the modeled CO abundance with the

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scaled NO_x emissions decreased by about 10% over Indonesia/Australia, for example, compared to the simulation with the a priori NO_x emissions. This reduction in CO represented about 20–30% of the contribution of emissions from the Indonesian/Australian region to the total abundance of CO and suggests that neglecting the influence of NO_x emissions (and of the emission of other precursors of O₃) on the CO chemistry could contribute to a significant bias in the CO source estimates. To accurately quantify the surface emissions of CO in an inverse modeling framework, it will be necessary to properly account for the chemical coupling of the CO-O₃-NO_x-hydrocarbon chemistry.

We found that although both TES and MOPITT provide global observational coverage, differences in the spatio-temporal sampling of the measurements contribute to differences between the a posteriori source estimates obtained with the two datasets. As discussed above, these differences were small (less than 20%) for most of the continental source regions considered here, but they are nonetheless significant. For North American emissions, to which there was less sensitivity in the inversion, the difference between the estimates from TES and MOPITT was about 75%, reflecting the influence of systematic errors in the analysis. It is likely that some of the discrepancies between previously published top-down emissions estimates could be due to the use of *ad hoc* state vectors based on geo-political boundaries, which are independent of the spatio-temporal resolution and precision of the data used in the analyses. Furthermore, the presence of systematic errors in the analysis will adversely impact the elements of the state vectors which are poorly constrained by the observations. We showed that although the a posteriori source estimates provided a significantly improved simulation of the TES and MOPITT data, regional residual biases remained in the simulated CO distribution, which are probably due to aggregation errors in the inversion state vector and to systematic models errors. The presence of these residuals, despite the consistency between the source estimates inferred from TES and MOPITT, suggests that reconciling the discrepancies between top-down source estimates will require properly characterizing systematic observation and model errors (chemical and transport) and conducting the inversion at a spatial-temporal resolution consistent with the information

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content of the observations. It also indicates the need to optimally combine boundary layer measurements of CO with TES and MOPITT data, along with observations of other tracers such as NO₂, which have similar sources as CO, in a multi-species inversion framework.

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Table 1. A priori and a posteriori source estimates.

Region	A priori ¹ (Tg CO/yr)				A posteriori TES Nov 4–16 (Tg CO/yr)	A posteriori MOPITT Nov 5–15 (Tg CO/yr)	A posteriori MOPITT Nov 5–28 (Tg CO/yr)	Difference between estimates from TES and MOPITT ³ (%)
	BB	BF	FF	Total				
North America	23	6	106	135	36	146	165	–75
Europe	8	17	85	110	132	111	111	19
Asia	102	93	171	367	511	531	483	–4
South America	69	19	25	113	118	141	157	–16
Northern Africa	99	21	19	139	131	119	174	10
Sub-equatorial Africa	78	10	6	95	173	184	185	–6
Indonesia/ Australia	49	7	13	69	155	185	165	–16
Rest of the World ²				1215	1390	1344	1336	3
Total				2243	2646	2761	2776	–4

¹ Sources represent emissions of CO from fossil fuel (FF) and biofuel combustion (BF) and biomass burning (BB), based on Duncan et al. (2003, 2007).

² The rest of the world source includes CO from the oxidation of methane and non-methane hydrocarbons.

³ Difference in TES a posteriori estimates relative to those from MOPITT calculated from data in early November.

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Table 2. Regional bias in the model simulation of CO¹.

Region	TES		MOPITT	
	A priori bias (%)	A posteriori bias (%)	A priori bias (%)	A posteriori bias (%)
Globe (60° S–0° N)	–12	0.1	–22	–0.8
Southern hemisphere (0°–5° S)	–18	–2	–27	–1
Southern Atlantic (35°–0° S, 40° W–5° E)	–18	–3	–27	–5
Southern Africa/Indian Ocean (35°–5° S, 15°–0° E)	–20	–5	–31	–7
Central Pacific Ocean (10° S–10° N, 180° W–0° W)	–14	0.3	–13	8
Northern hemisphere (25° N–60° N)	–7	1	–18	–3
North Pacific (25°–0° N, 175° W–120° W)	–10	–0.9	–24	–8
East Asia (25°–60° N, 110° E–135° E)	–4	3	–12	6

¹ The bias is calculated (model minus observations) with respect to CO columns from TES and MOPITT data, with the a priori and a posteriori emissions of CO in the model.

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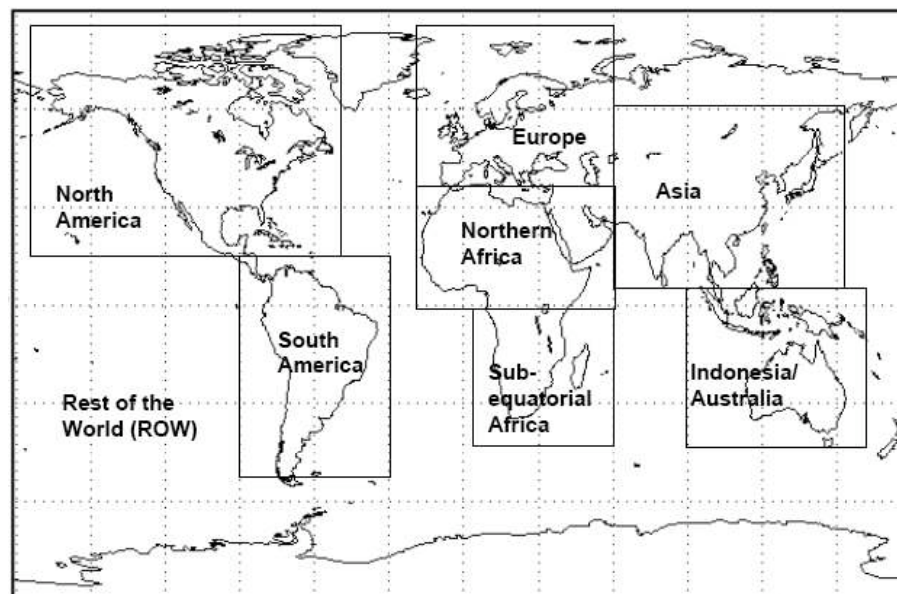


Fig. 1. Source regions comprising the state vector in the inversion analysis. The a priori estimates for emissions from these regions are listed in Table 1.

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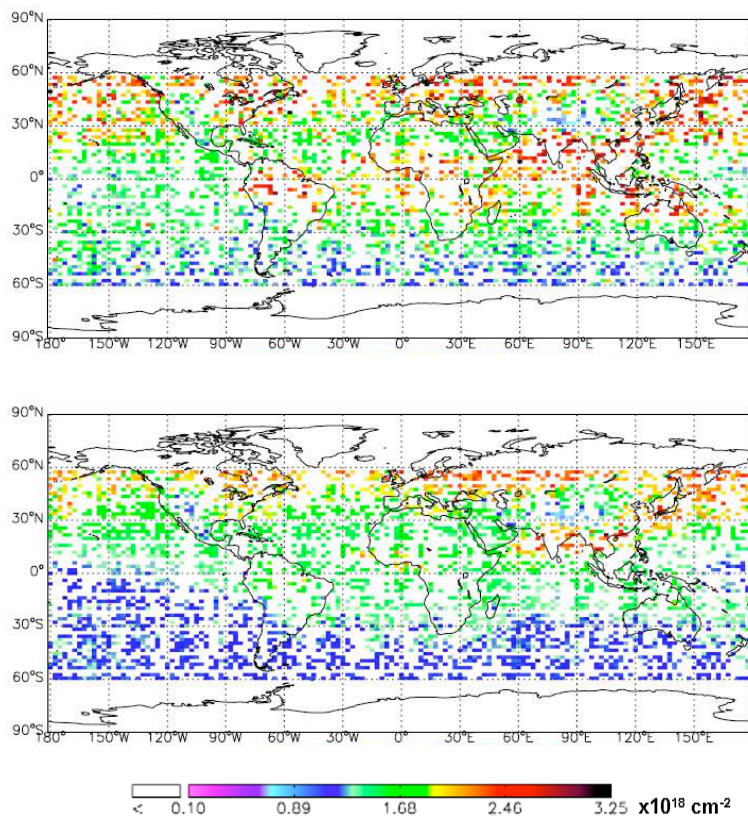


Fig. 2a, b. (a) Column abundances of CO (in units molecules cm^{-2}) from TES, averaged 4–15 November 2005. (b) Column abundances of CO from the GEOS-Chem model. The modeled fields were sampled along the TES orbit and smoothed using the TES averaging kernels and a priori profile. White areas are regions without observations.

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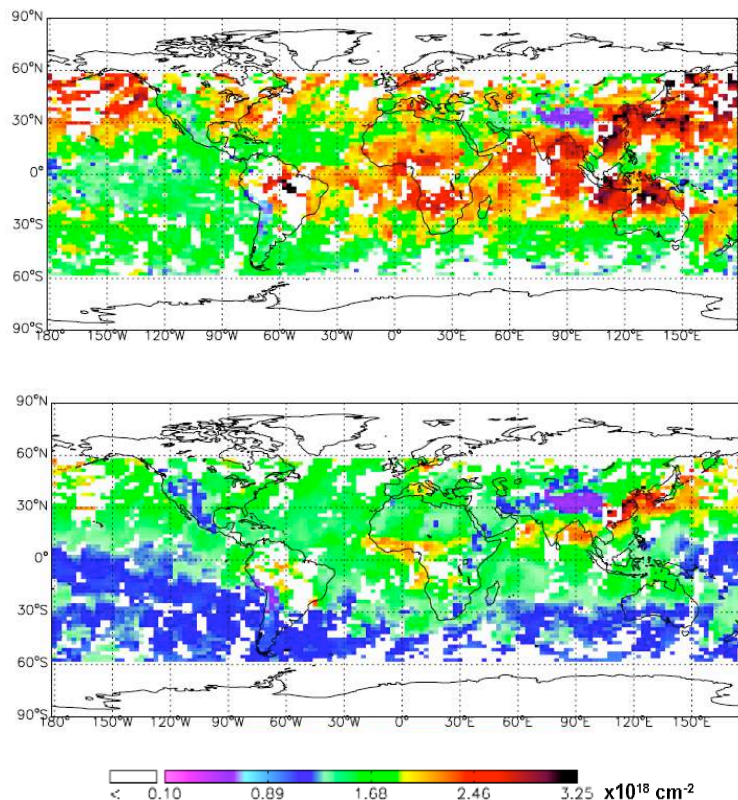


Fig. 2c, d. (c) Column abundances of CO (in units molecules cm^{-2}) from MOPITT, averaged 5–15 November 2004. (d) Column abundances of CO from the GEOS-Chem model. The modeled fields were sampled along the MOPITT orbit and smoothed using the MOPITT averaging kernels and a priori profile. White areas are regions without observations.

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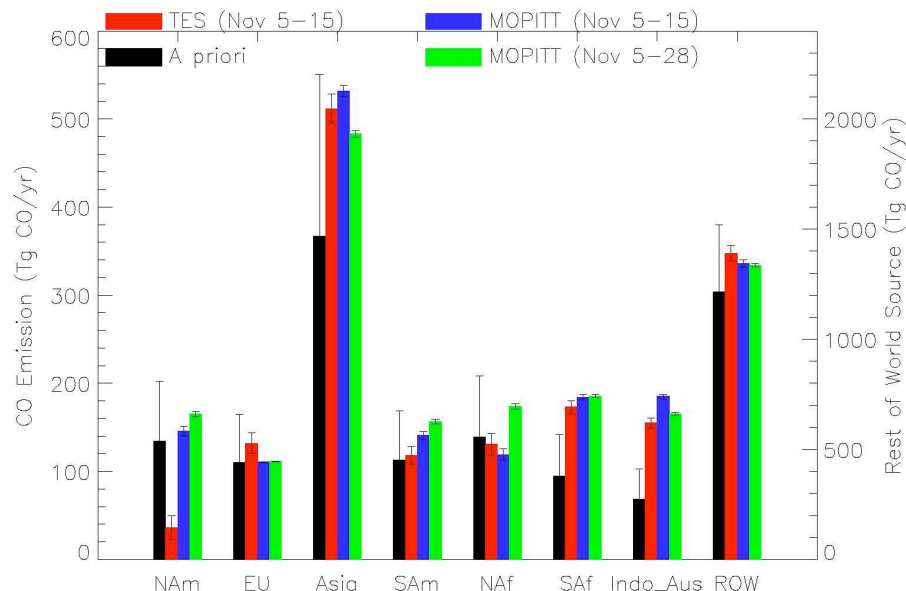


Fig. 3. A priori and a posteriori CO source estimates. Black bars indicate the a priori emission estimates, red bars are the a posteriori estimates inferred from TES data (4–15 November 2004), whereas the blue bars are the a posteriori estimates based on MOPITT data (5–15 November 2004). The green bars represent the a posteriori emission estimates obtained using MOPITT data for 5–28 November 2004. The regional definitions are: North America (NAM), Europe (EU), South America (SAM), North Africa (NAF), sub-equatorial Africa (SAF), Indonesia/Australia (Indo_Aus), and the rest of the world and the background chemical source of CO (ROW). Error bars indicate an uncertainty of $1\text{-}\sigma$.

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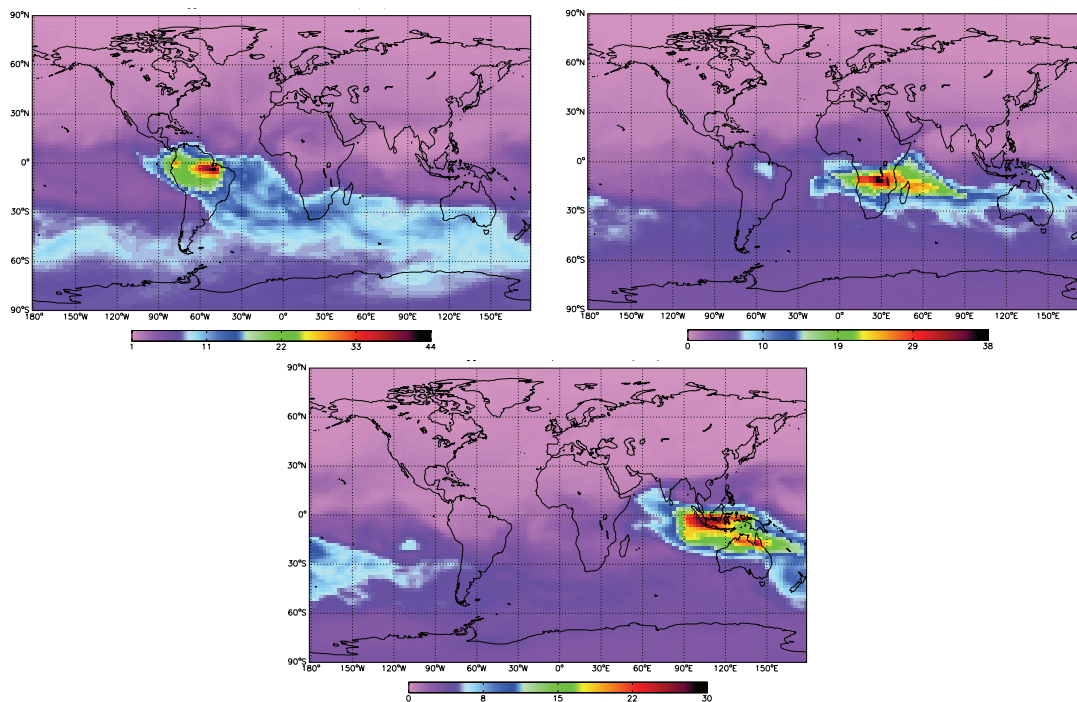


Fig. 4. Distribution of the tagged CO tracer for emissions of **(a)** South America, **(b)** sub-equatorial Africa, and **(c)** Indonesia/Australia. The tracer distributions were obtained using the a priori CO emissions and are shown for the upper troposphere at 8 km, averaged between 1–15 November 2004. Units are in ppb.

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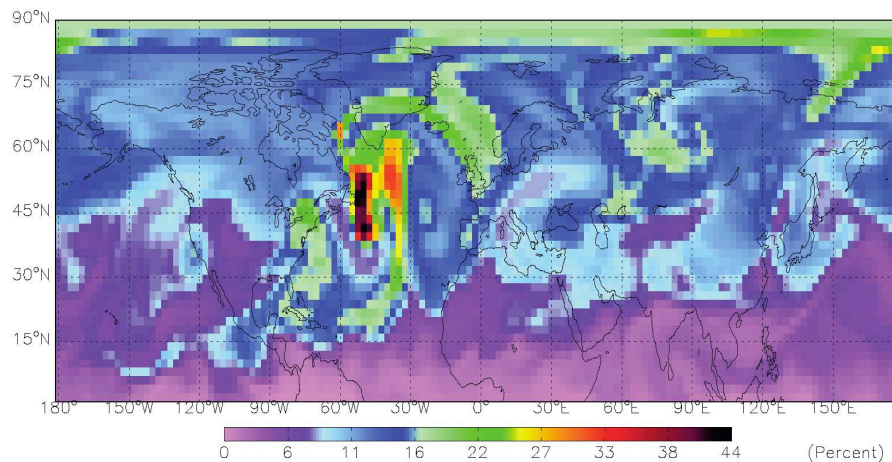


Fig. 5. Contribution of emissions of CO from North America (as defined in Fig. 1) to the total abundance of CO. The tracer distribution, as a percentage of total CO, is shown for 00:00 GMT on 5 November 2005 at about 5 km.

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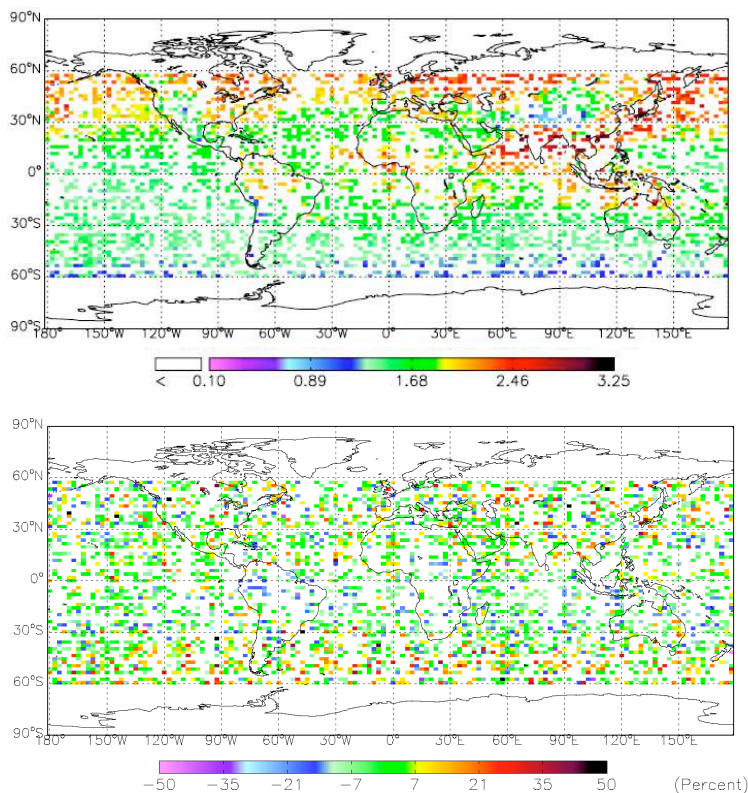


Fig. 6. (a) Column abundances of CO, averaged from 4–15 November 2004, from the GEOS-Chem simulation of the a posteriori emissions. Modeled fields transformed using the TES averaging kernels and a priori profiles. Units are 10^{18} molecules cm^{-2} . **(b)** The residuals expressed as a percent difference between the model and the TES observations (model minus TES). The TES data are shown in Fig. 2.

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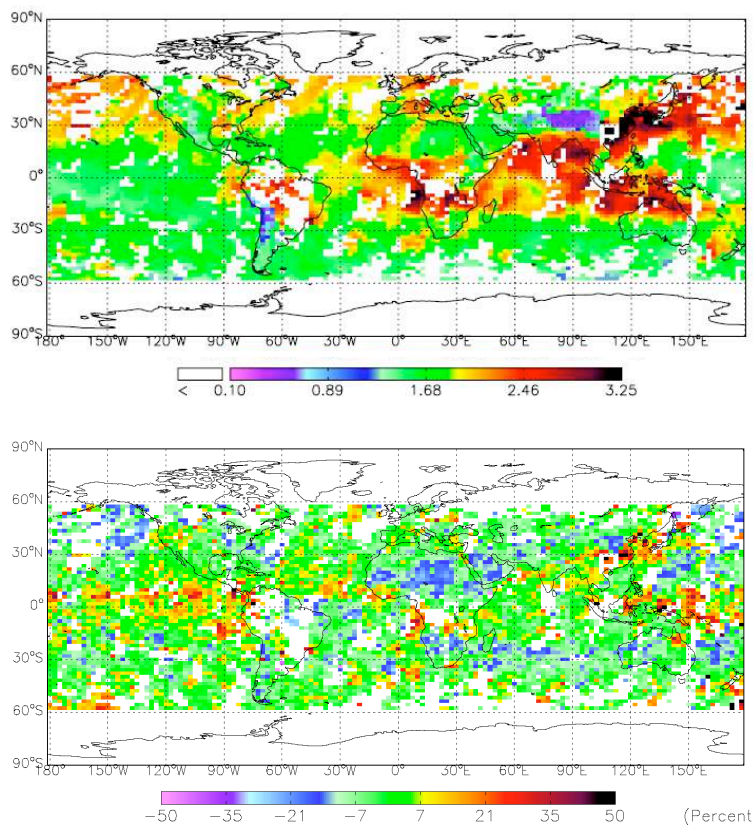


Fig. 7. (a) Column abundances of CO, averaged from 5–15 November 2004, from the GEOS-Chem simulation of the a posteriori emissions. Modeled fields transformed using the MOPITT averaging kernels and a priori profiles. Units are 10^{18} molecules cm^{-2} . **(b)** The residuals expressed as a percent difference between the model and the MOPITT observations (model minus MOPITT). The MOPITT data are shown in Fig. 2.

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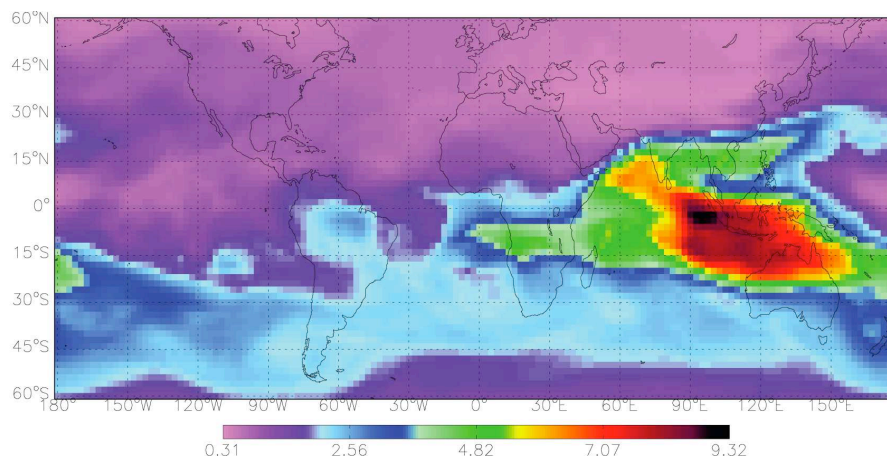


Fig. 8. Difference in modeled CO at 8 km in the GEOS-Chem model between simulations with the a priori emissions and with combustion related NO_x emissions scaled based on the regional scaling factors from the CO inversion. Shown are the a priori minus scaled NO_x simulations averaged between 1–15 November 2004. Units are in ppb.

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